

Application No.: 10/627,245
Amendment Dated: October 16, 2006
Reply to Office Action of May 15, 2006

Amendments to the Claims:

This listing of claims will replace all prior versions, and listings of claims in the application.

Listing of Claims:

1. (currently amended) A method for testing agents for effect on human cardiac cells comprising the steps of

~~culturing~~ deriving cardiomyocytes ~~derived~~ by *in vitro* culture from human embryonic stem cells;

piercing a single cardiomyocyte with an electrode so that the transmembrane action membrane of that cardiomyocyte can be electrically measured;

measuring the transmembrane action potential of ~~at least one~~ the single cardiomyocyte;

assessing the transmembrane action potential of the cardiomyocyte to characterize the cardiomyocyte as to the cell type of the human heart that the action potential most resembles among the cell types selected from the group consisting of ventricular, atrial and nodal cell types;

exposing the cardiomyocyte to the agent; and

observing whether the action potential of the cardiomyocyte changes after the exposure to the agent.

2. (cancelled).

3. (currently amended) The method of claim 1 wherein the ~~culturing~~ deriving is conducted by permitting the human embryonic stem cells to form embryoid bodies and wherein the measuring includes impaling the single cardiomyocyte within an embryoid body with ~~an~~ the electrode.

4-6. (cancelled).

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7. (currently amended) A method for testing agents for their effect on the electrical properties of the HERG channel in human cardiac cells comprising the steps of
~~culturing~~ deriving cardiomyocytes ~~derived~~ by *in vitro* culture from human embryonic stem cells;

inserting an electrode into the interior of a single ~~at least one~~ cardiomyocyte in culture
in order to be able to measure the transmembrane action potential of the cardiomyocyte;

measuring the duration of the transmembrane action potential of the cardiomyocyte;

assessing the transmembrane action potential of the cardiomyocyte to characterize the cardiomyocyte as to the cell type of the human heart that the action potential most resembles among the cell types selected from the group consisting of ventricular, atrial and nodal cell types;

exposing the cardiomyocyte to the agent; and

observing whether the action potential duration is changed by the agent, as would be the case if the HERG channel is altered.

8. (cancelled).

9. (original) The method of claim 7 wherein the culturing is conducted by permitting the human embryonic stem cells to form embryoid bodies and wherein the measuring includes impaling an embryoid body with an electrode.

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10. (currently amended) A method for testing agents for their likelihood of triggering delayed after depolarization events in human cardiac cells comprising the steps of
~~culturing~~ deriving cardiomyocytes ~~derived~~ by *in vitro* culture from human embryonic stem cells;

inserting an electrode into the interior of a single cardiomyocyte in culture in order to be able to measure the transmembrane action potential of the cardiomyocyte;

obtaining a chart of the transmembrane action potential of the cardiomyocyte over time;

assessing the transmembrane action potential of the cardiomyocyte to characterize the cardiomyocyte as to the cell type of the human heart that the action potential most resembles among the cell types selected from the group consisting of ventricular, atrial and nodal cell types;

exposing the cardiomyocyte to the agent; and

observing whether a delayed after polarization event is triggered by the agent.

11. (cancelled).

12. (original) The method of claim 10 wherein the culturing is conducted by permitting the human embryonic stem cells to form embryoid bodies and wherein the measuring includes impaling an embryoid body with an electrode.

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13. (currently amended) A method for testing agents for their likelihood of triggering long QT syndrome in patients by testing human cardiac cells comprising the steps of ~~culturing deriving~~ cardiomyocytes ~~derived~~ by *in vitro* culture from human embryonic stem cells;

inserting an electrode into the interior of several single cardiomyocytes in the culture in order to be able to measure the transmembrane action potential of the cardiomyocytes;

obtaining a chart of the transmembrane action potential of a plurality of the ~~eardiomyocyte~~ cardiomyocytes over time;

assessing the transmembrane action potential of the cardiomyocytes to characterize the cardiomyocytes as to the cell type of the human heart that the action potential most resembles among the cell types selected from the group consisting of ventricular, atrial and nodal cell types;

exposing the ~~eardiomyocyte~~ cardiomyocytes to the agent; and

observing whether action potential duration is prolonged as an indicator of the risk of long QT syndrome by the agent in any of the cardiomyocytes.

14. (cancelled).

15. (original) The method of claim 13 wherein the culturing is conducted by permitting the human embryonic stem cells to form embryoid bodies and wherein the measuring includes impaling embryoid bodies with an electrode.

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16. (new) A method for testing agents for effect on human cardiac cells comprising the steps of

- culturing human embryonic stem cells by in vitro culture to produce embryoid bodies;
- selecting amongst the embryoid bodies for embryoid bodies which demonstrate the presence of cardiomyocytes;
- piercing the embryoid body to place a fine electrode inside a single cardiomyocyte within the embryoid body so that the transmembrane action membrane of that cardiomyocyte can be electrically measured;
- measuring the transmembrane action potential of the single cardiomyocyte;
- assessing the transmembrane action potential of the cardiomyocyte to characterize the single cardiomyocyte as to the cell type of the human heart that the action potential most resembles among the cell types selected from the group consisting of ventricular, atrial and nodal cell types;
- exposing the cardiomyocyte to the agent; and
- observing whether the action potential of the cardiomyocyte changes after the exposure to the agent.